#2 OIPE

RAW SEQUENCE LISTING DATE: 11/20/2001 PATENT APPLICATION: US/09/965,522 TIME: 10:10:19

Input Set : N:\Crf3\RULE60\09965522.txt
Output Set: N:\CRF3\11202001\1965522.raw

SEQUENCE LISTING

```
3 (1) GENERAL INFORMATION:
      5
             (i) APPLICANT: Lal, Preeti
                            Bandman, Olga
      6
            (ii) TITLE OF INVENTION: NOVEL HUMAN SODIUM-DEPENDENT
      8
      9
                                      PHOSPHATE CO-TRANSPORTER
           (iii) NUMBER OF SEQUENCES: 7
     11
            (iv) CORRESPONDENCE ADDRESS:
     13
                  (A) ADDRESSEE: Incyte Pharmaceuticals, Inc.
     14
     15
                  (B) STREET: 3174 Porter Drive
                                                               ENTERED
                  (C) CITY: Palo Alto
     16
     17
                  (D) STATE: CA
                  (E) COUNTRY: US
     18
     19
                  (F) ZIP: 94304
             (V) COMPUTER READABLE FORM:
     21
                  (A) MEDIUM TYPE: Diskette
     22
     23
                  (B) COMPUTER: IBM Compatible
     24
                  (C) OPERATING SYSTEM: DOS
     25
                  (D) SOFTWARE: FastSEQ Version 2.0
     27
            (vi) CURRENT APPLICATION DATA:
C--> 28
                  (A) APPLICATION NUMBER: US/09/965,522
C--> 29
                  (B) FILING DATE: 26-Sep-2001
     30
                  (C) CLASSIFICATION:
     32
           (vii) PRIOR APPLICATION DATA:
     33
                  (A) APPLICATION NUMBER: 09/391,958
     34
                  (B) FILING DATE: 1999-09-08
     37
          (viii) ATTORNEY/AGENT INFORMATION:
     38
                  (A) NAME: Billings, Lucy J.
     39
                  (B) REGISTRATION NUMBER: 36,749
     40
                  (C) REFERENCE/DOCKET NUMBER: PF-0221 US
     42
            (ix) TELECOMMUNICATION INFORMATION:
     43
                  (A) TELEPHONE: 415-855-0555
                  (B) TELEFAX: 415-845-4166
        (2) INFORMATION FOR SEQ ID NO: 1:
     47
     49
             (i) SEQUENCE CHARACTERISTICS:
     50
                  (A) LENGTH: 401 amino acids
                  (B) TYPE: amino acid
     51
     52
                  (C) STRANDEDNESS: single
     53
                  (D) TOPOLOGY: linear
     55
           (vii) IMMEDIATE SOURCE:
     56
                  (A) LIBRARY: BRAITUT02
     57
                  (B) CLONE: 754412
     59
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
        Met Gln Val Asp Glu Thr Leu Ile Pro Arg Lys Val Pro Ser Leu Cys
     61
                                              10
     62
                           5
     63
         Ser Ala Arg Tyr Gly Ile Ala Leu Val Leu His Phe Cys Asn Phe Thr
     64
                     20
                                          25
```



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65 66	Thr	Ile	Ala 35	Gln	Asn	Val	Ile	Met 40	Asn	Ile	Thr	Met	Val 45	Ala	Met	Val
67 68	Asn	Ser 50	Thr	Ser	Pro	Gln	Ser 55		Leu	Asn	Asp	Ser 60	Ser	Glu	Val	Leu
69 70	Pro 65	Val	Asp	Ser	Phe	Gly 70	Gly	Leu	Ser	Lys	Ala 75	Pro	Lys	Ser	Leu	Pro 80
71 72	Ala	Lys	Ser	Ser	Ile 85	Leu	Gly	Gly	Gln	Phe 90	Ala	Ile	Trp	Glu	Arg 95	Trp
. 73 74	Gly	Pro	Pro	Gln 100	Glu	Arg	Ser	Arg	Leu 105	Cys	Ser	Ile	Ala	Leu 110	Ser	Gly
75 76	Met	Leu	Leu 115	Gly	Cys	Phe	Thr	Ala 120	Ile	Leu	Ile	Gly	Gly 125	Phe	Ile	Ser
77 78	Glu	Thr 130	Leu	Gly	Trp	Pro	Phe 135	Val	Phe	Tyr	Ile	Phe 140	Gly	Gly	Val	Gly
79 80	Cys 145	Val	Cys	Cys	Leu	Leu 150	Trp	Phe	Val	Val	Ile 155	Tyr	Asp	Asp	Pro	Val 160
81 82	Ser	Tyr	Pro	Trp	Ile 165	Ser	Thr	Ser	Glu	Lys 170	Glu	Tyr	Ile	Ile	Ser 175	Ser
83 84	Leu	Lys	Gln	Gln 180	Val	Gly	Ser	Ser	Lys 185	Gln	Pro	Leu	Pro	Ile 190	Lys	Ala
85 86	Met	Leu	Arg 195	Ser	Leu	Pro	Ile	Trp 200	Ser	Ile	Cys	Leu	Gly 205	Cys	Phe	Ser
87 88	His	Gln 210	Trp	Leu	Val	Ser	Thr 215	Met	Val	Val	Tyr	Ile 220	Pro	Thr	Tyr	Ile
89 90	Ser 225	Ser	Val	Tyr	His	Val 230	Asn	Ile	Arg	Asp	Asn 235	Gly	Leu	Leu	Ser	Ala 240
91 92	Leu	Pro	Phe	Ile	Val 245	Ala	Trp	Val	Ile	Gly 250	Met	Val	Gly	Gly	Tyr 255	Leu
93 94	Ala	Asp	Phe	Leu 260	Leu	Thr	Lys	Lys	Phe 265	Arg	Leu	Ile	Thr	V al 270	Arg	Lys
95 96			275			_		280					Leu 285			
97 98		290					295					300	Ala			
99 100	Leu 30!		Cys	Gly	Leu	Ser 310		Leu	Cys	Gln	Ser	_	Ile	Tyr	Ile	Asn 320
101			ı Ası	, Ile	ala			у Туі	r Sei	r Se	r Ph	e Lei	ı Met	: Gl	y Ala	a Ser
102					325					33					33	
103	Arg										e Vai	l Pro	Thi			r Gly
104				340							_	_	_	350		
105	Phe	e Leu			Glr	n Asr	Pro			e Gl	y Tr	o Arc			L Phe	e Phe
106	_	_	355			_	_	360		_	_		365			
107	Let			≥ Ala	a val	L Asr			ı GIŞ	y Lei	u Lei			: ье	1 116	e Phe
108	01 -	370					375				- 01.	380		. .	. mъ.	
109			1 Ale	a ASI	y va.			ı Tr	O AL	я гъ			у гул	з це	ı TII.	r Arg
110	38					390	,				39	,				400
111	Let		י אשם ר	PTON	EOP.	SEC	י חד	٠								
114 116	(2)	INFO		QUENC												
110		(± ,	, 511	50 TH	-L CI		- 1 LI L \ .									



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```
117
             (A) LENGTH: 1643 base pairs
118
             (B) TYPE: nucleic acid
119
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
120
122
       (vii) IMMEDIATE SOURCE:
123
             (A) LIBRARY: BRAITUT02
124
              (B) CLONE: 754412
126
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
    AGAACGGTGA GGATGACCGA CGTATAGGCG AGAGCCTAGG TACGCCATGC CAGGTCACCG
128
                                                                          60
    GTCCGGCAAT TCCCGGGTCG ACCCACGCGT CCGCTTGGAG GGACGCTGGG TTCAACTTGA
129
                                                                         120
130
    180
    GCCACCAAGA CAGAGTTGAG TCCCACAGCA AGGAGAGCA AGAACGCACA AGATATGCAA
131
                                                                         240
132
    GTGGATGAGA CACTGATCCC CAGGAAAGTT CCAAGTTTAT GTTCTGCTCG CTATGGAATA
                                                                         300
133
    GCCCTCGTCT TACATTTCTG CAATTTCACA ACGATAGCAC AAAATGTCAT CATGAACATC
                                                                         360
    ACCATGGTAG CCATGGTCAA CAGCACAAGC CCTCAATCCC AGCTCAATGA TTCCTCTGAG
134
                                                                         420
    GTGCTGCCTG TTGACTCATT TGGTGGCCTA AGTAAAGCCC CAAAGAGTCT TCCTGCAAAG
135
                                                                         480
    TCCTCAATAC TTGGGGGTCA GTTTGCAATT TGGGAAAGGT GGGGCCCTCC ACAAGAACGA
136
                                                                         540
137
    AGCAGACTCT GCAGCATTGC TTTATCAGGA ATGTTACTGG GATGCTTTAC TGCCATCCTC
                                                                         600
    ATAGGTGGCT TCATTAGTGA AACCCTTGGG TGGCCCTTTG TCTTCTATAT CTTTGGAGGT
138
                                                                         660
    GTTGGCTGTG TCTGCTGCCT TCTCTGGTTT GTTGTGATTT ATGATGACCC CGTTTCCTAT
    CCATGGATAA GCACCTCAGA AAAAGAATAC ATCATATCCT CCTTGAAACA ACAGGTCGGG
                                                                         780
140
    TCTTCTAAGC AGCCTCTTCC CATCAAAGCT ATGCTCAGAT CTCTACCCAT TTGGTCCATA
141
                                                                         840
    TGTTTAGGCT GTTTCAGCCA TCAATGGTTA GTTAGCACAA TGGTTGTATA CATACCAACT
142
                                                                         900
    TACATCAGCT CTGTGTACCA TGTTAACATC AGAGACAATG GACTTCTATC TGCCCTTCCT
143
                                                                         960
    TTTATTGTTG CCTGGGTCAT AGGCATGGTG GGAGGCTATC TGGCAGATTT CCTTCTAACC
144
                                                                        1020
145
    AAAAAGTTTA GACTCATCAC TGTGAGGAAA ATTGCCACAA TTTTAGGAAG TCTCCCCTCT
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146
    TCAGCACTCA TTGTGTCTCT GCCTTACCTC AATTCCGGCT ATATCACAGC AACTGCCTTG
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    CTGACGCTCT CTTGCGGATT AAGCACATTG TGTCAGTCAG GGATTTATAT CAATGTCTTA
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    GATATTGCTC CAAGGTATTC CAGTTTTCTC ATGGGAGCAT CAAGAGGATT TTCGAGCATA
148
                                                                        1260
    GCACCTGTCA TTGTACCCAC TGTCAGCGGA TTTCTTCTTA GTCAGGACCC TGAGTTTGGG
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149
150
    TGGAGGAATG TCTTCTTCTT GCTGTTTGCC GTTAACCTGT TAGGACTACT CTTCTACCTC
                                                                        1380
151
    ATATTTGGAG AAGCAGATGT CCAAGAATGG GCTAAAGAG GAAAACTCAC TCGTTTATGA
                                                                        1440
    AGTTATCCCA CCTTGGATGG AAAAGTCATT AGGCACCGTA TTGCATAAAA TAGAAGGCTT
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                                                                        1500
    CCGTGATGAA AATACCAGTG AAAAGATTTT TTTTTCCTGT GGCTCTTTTC AATTATGAGA
                                                                        1560
154
    TCAGTTCATT ATTTTATTCA GACTTTTTTT TGAGAGAAAT GTAAGATGAA TAAAAATTCA
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                                                                        1643
157 (2) INFORMATION FOR SEQ ID NO: 3:
159
         (i) SEQUENCE CHARACTERISTICS:
160
             (A) LENGTH: 467 amino acids
161
             (B) TYPE: amino acid
162
             (C) STRANDEDNESS: single
163
             (D) TOPOLOGY: linear
165
       (vii) IMMEDIATE SOURCE:
166
             (A) LIBRARY: GenBank
167
             (B) CLONE: 450532
169
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
171
    Met Gln Met Asp Asn Arg Leu Pro Pro Lys Lys Val Pro Gly Phe Cys
172
                                        10
173
    Ser Phe Arg Tyr Gly Leu Ser Phe Leu Val His Cys Cys Asn Val Ile
```



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174				20					25					30		
175	Ile	Thr	Ala	Gln	Arg	Ala	Cys	Leu	Asn	Leu	Thr	Met	Val	Val	Met	Val
176			35					40					45			
177	Asn	Ser	Thr	Asp	Pro	His	Gly	Leu	Pro	Asn	Thr	Ser	Thr	Lys	Lys	Leu
178		50					55		•			60				
179	Leu	Asp	Asn	Ile	Lys	Asn	Pro	Met	\mathtt{Tyr}	Asn	Trp	Ser	Pro	Asp	Ile	Gln
180	65					70					75					80
181	Gly	Ile	Ile	Leu	Ser	Ser	Thr	Ser	${ t Tyr}$	Gly	Val	Ile	Ile	Ile		Val
182					85					90					95	
183	Pro	Val	Gly	_	Phe	Ser	Gly	Ile	_	Ser	Thr	Lys	Lys		Ile	Gly
184		_ •	_	100	_	_	_		105	_	_	_		110		
185	Phe	Ala		Cys	Leu	Ser	Ser		Leu	Ser	Leu	Leu		Pro	Pro	Ala
186			115	<u></u>			_	120	1		_	_	125		~ 3	a 1
187	АТА	_	тте	GLY	val	АТа	Trp	vaı	vaı	vaı	Cys		Ala	val	GIn	GIĀ
188	31 0	130	C1 -	c1	, Tla	17 a 1	135	mh se	31-	C1 =	Dha	140	T1.	M	17a 1	T
189		Ата	GIN	GTA	ıте		Ala	Thr	Ala	GIN		GIU	тте	туг	val	LуS 160
190	145	31.	Dwo	D===	T 0	150	Arg	c1	A '	т о	155	Com	Wat	C ~ ~	mh ~	
191 192	тгр	Ald	PIO	PLO	165	GIU	Arg	СТА	Arg	170	THE	ser	Mec	ser	175	ser
193	Clv	Dho	Tau	Ten		Dro	Phe	Tla	Val		Leu	Va l	Thr	G1 17		Tlo
194	GIY	FILE	Бец	180	GLY	110	FIIC	116	185	пец	пеп	Val	1111	190	Val	110
195	Cvs	Glu	Ser		Glv	Trn	Pro	Met		Phe	Tvr	Tle	Phe		Δla	Cvs
196	CYB	Olu	195	шеч	017		110	200	141	1110	-1-		205	011		0,10
197	Glv	Cvs		Va 1	Cvs	Leu	Leu		Phe	Val	Leu	Phe		Asp	Asp	Pro
198	0-1	210			-1-		215					220	-1-	F		
199	Lvs	Asp	His	Pro	Cvs	Ile	Ser	Ile	Ser	Glu	Lvs	Glu	Tvr	Ile	Thr	Ser
200	225	-			-	230					235		-			240
201	Ser	Leu	Val	Gln	Gln	Val	Ser	Ser	Ser	Arg	Gln	Ser	Leu	Pro	Ile	Lys
202					245					250					255	
203	Ala	Ile	Leu	Lys	Ser	Leu	Pro	Val	Trp	Ala	Ile	Ser	Ile	Gly	Ser	Phe
204				260					265					270		
205	Thr	Phe		${\tt Trp}$	Ser	His	Asn	Ile	Met	Thr	Leu	Tyr	Thr	Pro	Met	Phe
206			275					280					285			
207	Ile		Ser	Met	Leu	His	Val	Asn	Ile	Lys	Glu		Gly	Phe	Leu	Ser
208		290					295				_	300		_		
209		Leu	Pro	Tyr	Leu		Ala	Trp	Ile	Cys		Asn	Leu	Ala	Gly	
210	305	_			_,	310		_	_		315	_				320
211	Leu	Ser	Asp	Pne		Leu	Thr	Arg	Asn		Leu	Ser	vaı	шe		val
212	3	T	T	nh -	325	31.	31.	~1	Dh.	330	T	D	7.7.	T1_	335	61
213	Arg	гуѕ	ьeu		Thr	Ala	Ala	GIY		Leu	Leu	Pro	Ala		Pne	GIY
214 215	Wa I	Crra	T 011	340	П•••	T 011	Com	Com	345	Dho	M	Com	т1 о	350	Tlo	Dho
216	val	Cys	355	PIO	TÄT	ьеu	Ser	360	1111	Pne	TAT	ser	365	Val	TTE	PHE
217	T.011	Tla		בוג	Gl v	λla	Thr		Sor	Dha	Cve	T.au		G1 v	V = 1	Dhe
218	БСи	370	Бец	AIG	GLY	AIG	375	GIY	361	rne	Cys	380	GLY	GLY	Val	rnc
219	Tle		Glv	Len	Asp	Tle	Ala	Pro	Ara	Ͳϒϒ	Phe		Phe	Tle	Tivs	Δla
220	385		1			390			*** 9	-1-	395	- I			2,5	400
221		Ser	Thr	Leu	Thr		Met	Ile	Glv	Glv		Ile	Ala	Ser	Thr	
222	4 -				405	- 4			2	410					415	
										-						



RAW SEQUENCE LISTING
PATENT APPLICATION: US/09/965,522

DATE: 11/20/2001 TIME: 10:10:19

223 224	Thr G	Sly	Leu	Ile 420	Leu	Lys	Gln	Asp	Pro 425	Glu	Ser	Ala	Trp	Phe 430	Lys	Thr
225 226	Phe I	le	Leu 435	Met	Ala	Ala	Ile	Asn 440	Val	Thr	Gly	Leu	Ile 445	Phe	Tyr	Leu
227	Ile V	/al		Thr	Ala	Glu	Ile		Asp	Trp	Ala	Lys	Glu	Lys	Gln	His
228	4	50					455					460				
229	Thr A	rg	Leu													
230	465															
232	(2) IN															
234	(i)				ARACI										
235			(A) LENGTH: 560 amino acids													
236			(B) TYPE: amino acid													
237			(C) STRANDEDNESS: single (D) TOPOLOGY: linear													
238 240	/ ***					OURCE		4.1			,.					
240	(^ 1					: Ge		ak .								
242						5074		11								
244	(x	ri)						N: SI	EQ II	NO:	4:					
246	Met G	•										Ala	Glv	Arσ	Ala	Leu
247	1			5	5					10			_	,	15	
248	Gly A	ırg	Leu	His	Arg	Leu	Leu	Glu	Lys	Arg	Gln	Glu	Gly	Ala	Glu	Thr
249	_	_		20	-				25	_				30		
250	Leu G	lu	Leu	Ser	Ala	Asp	Gly	Arg	Pro	Val	Thr	Thr	His	Thr	Arg	Asp
251			35					40					45			
252	Pro P	ro	Val	Val	Asp	Cys	Thr	Cys	Phe	Gly	Leu	Pro	Arg	Arg	Tyr	Ile
253		50					55					60				
254	Ile A	Ala	Ile	Met	Ser		Leu	Gly	Phe	Cys		Ser	Phe	Gly	Ile	
255	65			_		70			_		75	_	_	_		80
256	Cys A	Asn	Leu	Gly		Ala	Ile	Val	Ser		Val	Asn	Asn	Ser		Thr
257	77.2 A		61	61	85	17-1	37 <u>~</u> 1	37- 1	C1	90	71-	C1 =	Dho	7 ~ ~	95	7 0 0
258 259	His A	arg	GTA	100	HIS	val	val	Val	105	гуу	Ald	GIII	Pile	110	ттр	Asp
260	Pro G	27 11	Thr		Glv	T.e.11	Tle	Hic		Ser	Phe	Dhe	Tro		Ψvr	Tle
261	110 0	Lu	115	Val	GLY	пси	110	120	OLY	001	1110	1110	125	011	-1-	110
262	Val T	hr		Tle	Pro	Glv	Glv		Tle	Cvs	Gln	Lvs		Ala	Ala	Asn
263		.30	01			1	135			-1-		140				
264	Arg V	/al	Phe	Gly	Phe	Ala	Ile	Val	Ala	Thr	Ser	Thr	Leu	Asn	Met	Leu
265	145			_		150					155					160
266	Ile P	Pro	Ser	Ala	Ala	Arg	Val	His	Tyr	Gly	Cys	Val	Ile	Phe	Val	Arg
267					165					170					175	
268	Ile I	ieu	Gln	Gly	Leu	Val	Glu	Gly	Val	Thr	Tyr	Pro	Ala	Cys	His	Gly
269				180					185					190		
270	Ile T	rp.	Ser	Lys	Trp	Ala	Pro		Leu	Glu	Arg	Ser	Arg	Leu	Ala	Thr
271			195					200					205			
272	Thr A		Phe	Cys	Gly	Ser		Ala	Gly	Ala	Val		Ala	Met	Pro	Leu
273		210	_	_			215	_		_	_	220		_,		
274	Ala G	Sly	Val	Leu	Val		Tyr	Ser	Gly	Trp		Ser	Val	Phe	Tyr	
275	225	11	G	D1	~ 1	230	DI: -	m	m	* `	235	m	T = ···	T =	37.5 3	240
276	Tyr G	тА	ser	rne	GТĀ	тте	Pne	тrр	тyr	ьeu	rne	Trp	ьeu	ьeu	vaı	ser



VERIFICATION SUMMARY

PATENT APPLICATION: US/09/965,522

DATE: 11/20/2001 TIME: 10:10:20

Input Set : N:\Crf3\RULE60\09965522.txt
Output Set: N:\CRF3\11202001\1965522.raw

L:28 M:220 C: Keyword misspelled or invalid format, [(A) APPLICATION NUMBER:]

L:29 M:220 C: Keyword misspelled or invalid format, [(B) FILING DATE:]